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Diversity and catalytic potential of PAH-specific ring-hydroxylating dioxygenases from a hydrocarbon-contaminated soil

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Abstract

Ring-hydroxylating dioxygenases (RHDs) catalyze the initial oxidation step of a range of aromatic hydrocarbons including polycyclic aromatic hydrocarbons (PAHs). As such, they play a key role in the bacterial degradation of these pollutants in soil. Several PCR-based methods have been implemented to assess the diversity of RHDs in soil, allowing limited sequence-based predictions on RHD function. In the present study, we developed a method for the isolation of PAH-specific RHD gene sequences of Gram-negative bacteria, and for analysis of their catalytic function. The genomic DNA of soil PAH degraders was labeled *in situ* by stable isotope probing, then used to PCR amplify sequences specifying the catalytic domain of RHDs. Sequences obtained fell into five clusters phylogenetically linked to RHDs from either Sphingomonadales or Burkholderiales. However, two clusters comprised sequences distantly related to known RHDs. Some of these sequences were cloned in-frame in place of the corresponding region of the *phnAla* gene from *Sphingomonas* CHY-1 to generate hybrid genes, which were expressed in *E. coli* as chimerical enzyme complexes. Some of the RHD chimeras were found to be competent in the oxidation of 2- and 3-ring PAHs, but other appeared unstable. Our data are interpreted in structural terms based on 3D-modeling of the catalytic subunit of hybrid RHDs. The strategy described herein might be useful for exploring the catalytic potential of the soil metagenome and recruit RHDs with new activities from uncultured soil bacteria.

Keywords : Dioxygenases ; metagenomic DNA ; catalytic domain ; hybrid enzymes ; bioremediation

1 Introduction

2
3 Ring-hydroxylating dioxygenases (RHDs) are bacterial metalloenzymes that catalyze the first
4 step in the biodegradation of a variety of aromatic hydrocarbons. Some of them are able to
5 oxidize highly resistant compounds such as polycyclic aromatic hydrocarbons (PAHs)
6 (Jouanneau et al. 2011; Jouanneau et al. 2006). RHDs are two or three-component systems,
7 consisting of an oxygenase which bears the enzyme active site, and specific electron carriers
8 which deliver reductant to the oxygenase for oxygen activation during catalysis (Parales and
9 Resnick 2006). The oxygenase component is most often an heterohexamer $\alpha_3\beta_3$. Structural
10 studies on representative oxygenases showed that the α subunit is composed of two domains,
11 a Rieske domain that binds a [2Fe-2S] cluster, and the catalytic domain containing the
12 substrate-binding site and a mononuclear Fe(II) center where oxygenation of the substrate
13 takes place (Ferraro et al. 2005). Bacterial RHDs form a large enzyme family that can be
14 divided into four to five subgroups based on phylogenetic comparisons of their α subunit
15 amino acid sequences (Kweon et al. 2008; Nam et al. 2001). In this respect, PAH-specific
16 RHDs fall into two clusters of enzymes, those found in Gram-positive bacteria and those
17 found in Gram-negative bacteria (Jouanneau et al. 2011).

18 While the classification of RHDs essentially relies on enzymes found in cultured bacteria,
19 there is compelling evidence that bacterial species recalcitrant to cultivation outnumber
20 cultivable ones by several orders of magnitude in soils and sediments (Cole et al. 2010).
21 Hence, the biodiversity of bacterial enzymes in soil, including RHDs, is certainly much
22 greater than that considered in current classifications, as exemplified by a recent study of
23 dioxygenase genes by a metagenomics approach (Iwai et al. 2010).

24 The diversity of RHDs associated to PAH-polluted sites has been investigated by PCR-based
25 analysis of sequences encoding a portion of the α subunit gene. In marine sediments, RHD

1 genes appeared to be mainly represented by sequences related to the *Cycloclasticus phnAI*
2 gene (Lozada et al. 2008), whereas in marine microbial mats subjected to oil contamination,
3 gene sequences similar to *nahAc* found in Pseudomonads and to *nagAc* and *pahAc* found in
4 Betaproteobacteria were mostly detected (Bordenave et al. 2008). A variety of gene sequences
5 related to RHDs from Gram-negative bacteria have been found in soils and sediments
6 polluted by PAHs (Gomes et al. 2007; Ni Chadhain et al. 2006; Park and Crowley 2006; Zhou
7 et al. 2006). As a means to estimate the relative abundance of these genes in soil, quantitative
8 methods based on qPCR have been proposed (Cebon et al. 2008; Ding et al. 2010).
9 Differences in RHD biodiversity observed on PAH-polluted sites might reflect natural
10 variability, but they could also arise from the variety of primer sets used to PCR amplify
11 different portions of the α subunit genes (Iwai et al. 2011).

12 In a recent study, we explored the bacterial diversity of a PAH-contaminated site by stable
13 isotope probing, and found out that Betaproteobacteria were the main microorganisms
14 responsible for the degradation of the tricyclic hydrocarbon phenanthrene in soil (Martin et al.
15 2012). In order to examine the diversity of PAH-specific RHDs on the same experimental
16 site, ^{13}C -labeled DNA extracted from soil was used as a template to amplify a large segment
17 of the α subunit gene, extending over the catalytic domain. Phylogenetic analysis revealed
18 that the sequences obtained were related to RHDs found in Sphnigomonadales and
19 Burkholderiales. As a means to further investigate the enzymatic function associated to these
20 microbial sequences, some of them were inserted in place of a gene portion specifying the
21 catalytic domain of the RHD from *Sphingomonas* CHY-1 (Jakoncic et al. 2007b; Jouanneau
22 et al. 2006). Hybrid genes were expressed together with other components of strain CHY-1
23 enzyme in *E. coli*, and the activity of the resulting RHDs was assayed in the oxidation of 2-
24 and 3-ring PAHs. This study represents one of the first attempts to link partial sequences

- 1 retrieved from soil DNA to enzymatic function using a strategy based on exchange of
- 2 catalytic domains.
- 3

MATERIALS AND METHODS

Bacterial strains, plasmids and culture conditions

Bacterial strains and plasmids used in this study are listed in Tables 1 and 2. *E. coli* strains were grown in rich medium (Luria-Bertani) at 37°C with appropriate antibiotics as previously described (Demaneche et al. 2004).

Primer design and PCR amplification of sequences internal to RHD α subunit genes

Primers used in this study were designed after analysis of sequence alignments of RHD alpha subunits from representative and well established PAH degraders taken among Gram-negative bacteria. Sequences considered in the alignment are encoded by the following genes : *phnA1a* from *Sphingomonas* sp. CHY-1 (Demaneche et al. 2004), *bphA1f* from *Sphingobium yanoikuyae* B1 (Chadhain et al. 2007), *bphA1f* from *Novosphingobium aromaticivorans* (Romine et al. 1999), *phnA1f* from *Sphingomonas* sp. LH128 (Schuler et al. 2009), *phnA1* from *Cycloclasticus* sp. A5 (Kasai et al. 2003), the *phnAc* genes from *Burkholderia* sp. RP007 (Laurie and Lloyd-Jones 1999), *Acidovorax* sp. NA3 (Singleton et al. 2009), *Alcaligenes faecalis* AFK2 (accession # AB024945), *Delftia* sp. Cs1-4 (formerly *Burkholderia* Cs1-4; accession # AY367786), *Burkholderia* sp. DBT1 (Di Gregorio et al. 2004), and *ndoB* from *Pseudomonas putida* NCIB9816 (Simon et al. 1993). Two short, partially conserved sequences were selected for primer design, corresponding to FVCNYHGW and IGETSYR in PhnA1a from *Sphingomonas* sp. CHY-1. These sequences delimited a portion of the protein (residues 98-412) encompassing the almost entire catalytic domain (Jakoncic et al. 2007b).

One set of degenerate primers was intended to target a wide range of RHD genes from Proteobacteria. The nucleotide sequences of the forward (Phn-RHDf2) and reverse (Phn-

RHDR3) primers were TTCRYBTGCAAYTATCAYGGYTGG and CCYCKRTAVCWKGTYTCKCCRA, respectively. Another pair of primers, with a lower level of degeneracy, was used to specifically amplify sequences relevant to RHDs from Betaproteobacteria. The sequences of the forward and reverse primers were: TTCASYTGCACYTATCACGGCTGG (RHD-Beta-Grp1f) and CCKCKRTARGASGTYTCRCCAA (RHD-Beta-Grp1r). The template was a preparation of ¹³C-labeled DNA obtained through stable isotope probing. The DNA sample was extracted from a soil microcosm incubated for 5 days with 310 ppm ¹³C-phenanthrene, then purified by CsCl isopycnic ultracentrifugation as previously described (Martin et al. 2012). PCR was conducted with a Tpersonal thermocycler (Biometra) in a final volume of 50 µl consisting of 100 ng template DNA, 1× PCR buffer, 1.5 mM MgSO₄, 200 µM of each dNTP, 0.3 µM of each primer, 1 U of Kod Hot start DNA polymerase (Novagen). Amplification conditions were as follows: 95°C for 3 min, then 30 cycles of denaturation for 30 s at 95°C, annealing for 20 s and extension at 70°C for 16 s. Annealing was performed at 52°C for 10 cycles then at 50°C for 10 cycles and finally at 48°C for 10 cycles. When the RHD-Beta-Grp1f/r primer pair was used, PCR conditions were similar except that annealing was carried out at 63°C for 20 s and extension at 70°C for 15 s.

Construction of clone libraries and sequence analysis of amplicons

PCR products obtained with the Phn-RHDf2/R3 primer set were cloned into pSTBlue-1 using the perfectly blunt cloning kit (Novagen, Merck). Clones were screened for the occurrence of an insert of the correct size by PCR using the same primer set. A selection of positive clones was grown in a 96-well microtiter plate, resulting in a first library named DcatdioxA. A second library called DcatdioxB was obtained by cloning PCR products obtained with the RHD-Beta-Grp1f/r primers into pJET1.2 (Fermentas). The cloned inserts of

both libraries were subjected to nucleotide sequencing on both DNA strands (as performed by either Beckman Coulter genomics or Eurofins). The resulting sequences were translated, then aligned and compared using Clustal X. Phylogenetic analysis was performed by the neighbor-joining clustering method with bootstrap analysis, using the One-click software available on the www.phylogeny.fr website (Dereeper et al. 2008). The tree was formatted with Treedyn. Nucleotide sequences described in this study were deposited in the EMBL-EBI database under accession numbers HE795402-HE795451 (DcatdioxA library) and HE774402-HE774466 (DcatdioxB library).

Construction of hybrid RHD genes

Hybrid genes were constructed by replacing a portion of the CHY-1 *phnA1a* gene as follows. First, the *phnA1aA2a* pair of genes encoding PhnI was subcloned as a XbaI-HindIII fragment from pSD8 into pUC18 to give pFMA1. In this construction, the resulting PhnI α subunit carried a His-tag at the N-terminus. Plasmid pFMA1 was then used to replace a fragment internal to *phnA1a* by an appropriate sequence derived from the DcatdioxA clone library as outlined in supplementary figure S1. To this end, PCR fragments were amplified by using six cloned sequences, U3-18, U3-60, U3-72, U3-89, U3-112 and U3-116 as templates and primers introducing restriction sites for RsrII and AgeI at the 5'- and 3'-end, respectively. The fragments were digested by these two enzymes, and then cloned into corresponding sites of pFMA1 to generate hybrid genes showing a replacement of a 695 bp segment between nucleotide 545 and 1240 of the *phnA1a* coding sequence (Table 2). Two additional constructions were made using U3-16 and U3-124 as templates and primers introducing RsrII and MluI sites at the 5'- and 3'-end, respectively. The fragments were cloned into pFMA1, resulting in hybrid genes with a 580-bp sequence substitution between nucleotide 545 and 1125 (Table 2). All constructions were checked by nucleotide sequencing on both strands.

Overexpression and *in vivo* assays of RHD chimeras

Plasmid pFMA1 and vectors derived thereof carrying hybrid genes (Table 2) were introduced into strain JM109 together with plasmid pIBA34 by co-transformation. Plasmid pIBA34 carries the *phnA3* and *phnA4* genes encoding the ferredoxin and reductase components of CHY-1 RHD, respectively. The PhnA4 reductase was His-tagged at the N-terminus. Transformants were grown at 37°C until bacterial density reached 1.0 (OD₆₀₀), then induced with 1 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) and further incubated at 25°C for about 20 h. Cultures were centrifuged, washed and resuspended to a bacterial density of 2.0 (OD₆₀₀) in M9 medium containing 20 mM glucose. Bacterial suspensions were distributed in 2-ml Eppendorf tubes (1ml/tube), in which 0.5 μmole of PAH had been introduced as a solution in acetone. After solvent evaporation, bacteria were incubated on a rotary shaker at 25°C for 6 h. Tubes were centrifuged 2 min at 12,000 g, then 0.9 ml of supernatant medium was mixed with 9 μl of 2,3-dihydroxybiphenyl (0.1 or 1 mM; Sigma-Aldrich) as an internal standard. The aqueous fraction was extracted with an equal volume of ethyl acetate. Dried extracts were taken up in 0.2 ml acetonitrile and analyzed by GC/MS as *n*-butylboronate (NBB) derivatives as previously described (Schuler et al. 2008). Anthracene dihydrodiol was instead reacted with a silylation reagent (Silyl-991; Macherey-Nagel). Dihydrodiol derivatives were quantified on the basis of peak area using calibration curves generated with known amounts of naphthalene 1,2-dihydrodiol for NBB products, or anthracene 1,2-dihydrodiol for trimethylsilyl derivatives (Jouanneau et al. 2006). Salicylate hydroxylase activity was assayed as described previously (Demaneche et al. 2004). Briefly, induced cells were resuspended in 10 ml M9 medium as above, then incubated with 1 mM salicylate for 6 h. After pelleting bacteria by centrifugation and ethyl acetate extraction, catechol was detected by GC/MS as a NBB derivative.

Purification of his-tagged oxygenase components

The his-tagged PhnI oxygenase produced in JM109(pFMA1), as well as enzyme chimera produced from similar expression systems, were tentatively purified through a procedure inspired from that described previously (Jouanneau et al. 2006), with simplifications. Cultures of recombinant bacteria (0.8 L) were grown in 2-L flasks to an OD₆₀₀ of 1, then supplemented with 50 μ M Fe(NH₄)₂(SO₄)₂ and induced with 1 mM IPTG at 20°C for 20 h. Bacteria were harvested by centrifugation and kept frozen at -20°C. Frozen pellets were resuspended in twice the volume of PGE buffer consisting of 25 mM potassium phosphate pH 7.5, 0.5 M NaCl, 5% glycerol, 5% ethanol and 2 mM β -mercaptoethanol. The following steps were performed under anoxic conditions with argon-saturated buffers. After treatment with 0.5 mg/ml lysozyme for 20 min, suspensions were subjected to ultrasonication for 5 min on ice with a Vibra Cell apparatus operated in pulse mode (Fisher Bioblock Scientific, Illkirch, France). The lysates were centrifuged at 12,000 g for 20 min, and the resulting cell extracts were applied onto small columns (1.5 ml) of immobilized-Co²⁺ affinity chromatography resin (TALON, BD Biosciences Ozyme, France). The columns were washed successively with 15 ml of PGE buffer and 10 ml of PGE containing 10 mM imidazole. The proteins were eluted in approx. 1 ml of PGE lacking NaCl but containing 0.15 M imidazole and kept frozen in liquid nitrogen.

Biochemical analyses

Expression of recombinant RHDs in *E. coli* was analyzed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) on 10% slab gels using a SE 260 Mighty Small II system (Hoefer®, Inc). Gels were processed and stained as previously described (Jouanneau et al. 2006). For Western blot analysis, proteins were transferred to Hybond C

1 nitrocellulose membranes (Amersham Biosciences) in a Biometra apparatus as described
2 previously (Jouanneau et al. 1995). Membranes were processed for immunodetection of His-
3 tagged proteins using mice anti-Histag IgG coupled to horse radish peroxidase at a 1/3000
4 dilution ratio (Sigma-Aldrich). Blots were revealed using a luminescent reagent (ECL;
5 Amersham Biosciences) and a Fusion Fx7 imaging system (Vilbert Lourmat).
6 Protein concentrations were determined using the Bradford reagent with bovine serum
7 albumin as a standard.

8 **Molecular modeling**

10 The crystal structure of the oxygenase component (PhnI) of *Sphingomonas* CHY-1
11 RHD (PDB code: 2CKF) was used as a template to generate 3D-models for the hybrid
12 enzymes. To this end, all necessary residue replacements were simultaneously performed in
13 the relevant region of the PhnI catalytic domain (between amino acids 185 and 415 of the
14 α subunit), using COOT (Emsley et al. 2010). The backbone structure of the catalytic domain
15 was therefore maintained. The position of replaced amino acids was inspected and sometimes
16 adjusted to minimize interactions, using coordinates and electron density maps of the original
17 PhnI structure as guides. Water molecules were not modeled.

RESULTS

Isolation of RHD sequences from soil treated with ^{13}C -phenanthrene

In a previous study, we implemented a stable isotope probing approach to identify PAH degrading bacteria in a hydrocarbon-contaminated soil from a road runoff collecting facility (Martin et al. 2012). Analysis of a ^{13}C -labeled DNA fraction purified from soil incubated for 5 days with 310 ppm ^{13}C -phenanthrene, showed that phenanthrene degraders were dominated by Betaproteobacteria. Here, we made use of the same DNA fraction to investigate the biodiversity of RHD sequences possibly involved in PAH oxidation. To this end, we employed a PCR-based strategy to amplify a *ca* 940-bp internal fragment of the alpha subunit gene. The sequences, delimited by nucleotides 291 and 1234 in the *phnA1a* gene from strain CHY-1, extended between two partially conserved regions corresponding to the [2Fe-2S] cluster binding site on the 5'-end, and a region 130-bp from the 3'-end (Fig. S1). It covered part of the Rieske-cluster binding domain and the major part of the catalytic domain, including the ligands of the Fe(II) center at the active site and residues forming the substrate-binding pocket (Jakoncic et al. 2007a; Jakoncic et al. 2007b).

Primers used for PCR amplification were designed after alignment of RHD alpha subunits from representative PAH degraders. The Phn-RHDf2/R3 primer set, which showed a high degree of degeneracy, was intended to target all RHD α -subunit genes found in a selection of Proteobacteria (see Methods). PCR conditions affording specific amplification of the expected *phnA1a* fragment from strain CHY-1 genomic DNA were applied to recover RHD gene sequences from ^{13}C -labeled metagenomic soil DNA. From a total of 96 sequences analyzed, fifty were predicted to encode RHD catalytic domains of the expected size. Among the other clones, two contained RHD sequences bearing a stop codon, thirty-six contained sequences unrelated to RHDs, and eight yielded incomplete sequences.

Phylogenetic analysis of relevant translated products revealed that the RHD sequences were highly diverse and comparison with counterparts found in the databases showed that they fell into four clusters (Fig. 1). Cluster 1 featured 11 sequences sharing a high level of similarity (86-99% over 314 amino acids) with the α subunit of previously studied RHDs, including NahA1f from *Sphingomonas* LH128 (Schuler et al. 2009) and PhnA1a from *Sphingomonas* CHY-1 (Demaneche et al. 2004). Cluster 2 comprised 17 sequences, which appeared to be more distantly related to known RHDs, as similarity levels did not exceed 67% with the best matching α subunit (PhnA1a from strain CHY-1). Some of the sequences such as U3-89 best matched with PhnA1 from *Cycloclasticus* A5 (62%, (Kasai et al. 2003)). Cluster 3 contained 15 sequences showing less than 40% identity with relevant PAH-specific RHDs. On the other hand, this group was found to share significant sequence similarity with BphA1b from *Novosphingobium aromaticivorans* (87%; (Romine et al. 1999)) and AhdA1b from *Sphingobium* sp. P2 (84%; (Pinyakong et al. 2003b)). The latter proteins resembled the α subunit of oxygenases mainly found in Sphingomonadaceae, which do not catalyze dioxygenase reactions but instead function as salicylate hydroxylases (Jouanneau et al. 2007; Pinyakong et al. 2003b). Finally, 7 sequences were related to RHDs from Burkholderiales, four of which were closely similar and grouped with sequences in cluster 4 (see below). These sequences showed limited similarities with α subunits of known RHDs. In particular, none of the cloned sequences was related to the RHD from *Acidovorax* NA3 (Singleton et al. 2009), whereas bacteria belonging to this genus were found to be dominant among phenanthrene degraders identified by SIP in the soil of interest (Martin et al. 2012).

Given that PCR amplification from environmental DNA is often subjected to biases resulting in underrepresentation of some sequences in gene libraries, we used a second set of less degenerate primers designed to target the RHD α subunit genes from Betaproteobacteria. PCR amplicons obtained from the same sample of soil ^{13}C -DNA as template served to

1 generate a second library, from which 65 DNA sequences were analyzed. The deduced
2 protein sequences were all related to RHDs from Burkholderiales (Fig. 1), and could be
3 divided into two clusters. The majority of sequences belonged to cluster 4, and shared less
4 than 65% identity with the closest match in the UniProt database, the RHD α subunit from
5 *Burkholderia* sp. DBT1 (Di Gregorio et al. 2004). Interestingly, this strain initially isolated on
6 dibenzothiophene as sole carbon source was recently shown to degrade phenanthrene and
7 other PAHs (Andreolli et al. 2011), suggesting that its dioxygenase could oxidize these
8 hydrocarbons. The last cluster (cluster 5) consisted of 8 sequences highly similar to the α
9 subunit of *Acidovorax* NA3 RHD (>94% identity over 314 amino acids)(Fig. 1).

10 In order to assess the catalytic activity associated to some of the RHD sequences
11 described above, construction of enzyme chimeras was undertaken using the well-described
12 dioxygenase from *Sphingomonas* CHY-1 as a model.

14 **Construction of hybrid *phnA1a* genes and expression of RHD chimera in *E. coli***

15 In previous studies, we used a 12-kb plasmid (pSD9) for optimal expression of the
16 PhnI oxygenase from strain CHY-1 in either *E. coli* (Demaneche et al. 2004) or
17 *Pseudomonas putida* (Jouanneau et al. 2006). In order to facilitate the genetic constructions
18 described in this study (Table 2), a smaller expression plasmid was generated by cloning a
19 2.3-kb DNA fragment including the *phnA1a* and *phnA2a* structural genes in pUC18 under the
20 *lac* promoter (Fig. S1). The resulting plasmid (pFMA1) allowed consistent overproduction of
21 PhnI in *E. coli* JM109, and yielded RHD activity levels about 60% as high as the
22 BL21(DE3)(pSD9) expression system, when combined with plasmid pIBA34 expressing
23 accessory electron carriers PhnA4 and PhnA3 (data not shown).

24 Eight hybrid genes were constructed by exchange of a 695-bp fragment of *phnA1a* by
25 an equivalent fragment from sequences U3-18, U3-60, U3-72, U3-89, U3-112 and U3-116, or

1 by exchange of a 580-bp fragment for the U3-16 and U3-124 sequences (Table 2; see
2 Materials and Methods for details). Resulting plasmids were transferred into strain JM109
3 carrying pIBA34, and expression of RHD chimeras was examined. Analysis of whole cell
4 protein extracts from recombinant *E. coli* strains by Western blot using His-tag-specific
5 antibodies indicated that hybrid alpha subunits ($M_r \approx 52,000$) were produced in all cases (Fig.
6 2A). The second his-tagged protein detected with a M_r of 45,000 was the co-expressed
7 reductase PhnA4.

8 Three RHD chimera derived from the U3-18, U3-72 and U3-116 sequences were
9 found to be catalytically active and transformed naphthalene to the corresponding 1,2-
10 dihydrodiol at rates lower than the reference PhnI enzyme expressed under similar conditions
11 (Tables 2 and 3). All other constructions yielded no detectable dioxygenase activity. The
12 RHD derived from U3-18 and U3-116 also transformed phenanthrene, biphenyl and
13 anthracene at relative rates comparable to those found for PhnI (Table 3). The hybrid enzyme
14 made from the U3-72 sequence showed low but significant activity with phenanthrene and
15 biphenyl. Since the deduced amino acid sequences of U3-16 and U3-124 showed high
16 similarity with the catalytic domain of salicylate hydroxylases, we tested whether the relevant
17 hybrid enzymes would be competent in the hydroxylation of salicylate. No activity was
18 detected suggesting that hybrid enzymes were not stably produced in *E. coli*, as supported by
19 SDS-PAGE analysis of soluble extracts (data not shown).

20 Given that RHDs are multi-component enzymes, several reasons might explain the
21 observed poor activity of most of the studied chimeras, including wrong folding of the hybrid
22 alpha subunit, defective assembly of the $\alpha_3\beta_3$ hexamer or incorrect interaction of the hybrid
23 oxygenase component with the PhnA3 ferredoxin. We first examined whether the enzyme
24 constructions were produced in soluble form in the cytosolic fraction of *E. coli*. Immunoblot
25 analysis of such fractions indicated that only hybrid RHDs endowed with high activity (U3-18

and U3-116) were detected at levels comparable to PhnI (Fig. 2B). On the other hand, hybrid RHDs displaying little or no activity were barely detected in *E. coli* cytosol (Fig. 2B, lines 5-7). Since the His-tagged α -subunit of these hybrids was found in whole cell extracts (Fig. 2A), it is likely that they accumulated as inclusion bodies, suggesting that the lack of activity of those hybrid RHDs was primarily due to instability or misfolding.

We then tried to isolate hybrid RHDs by affinity chromatography (IMAC), taking advantage of the His-tag fused at the N-terminus of the α subunit. Only hybrids with high enough dioxygenase activity could be isolated in significant amount. The absorbance spectrum of the U3-116 hybrid isolated in this way was very similar to that of PhnI (Jouanneau et al. 2006), indicating that Fe-S centers were correctly inserted in the protein (data not shown). On the other hand, hybrid RHDs with no activity could not be isolated from *E. coli* soluble extracts suggesting that these constructions were essentially produced as insoluble proteins.

Structural insights based on 3D modeling of hybrid RHDs

To further analyze the structural consequences of the exchange of catalytic domain on hybrid RHDs, we took advantage of the known crystal structure of PhnI from strain CHY-1 to perform molecular modeling of the hybrids (Jakoncic et al. 2007b). The 4 hybrids derived from U3-18, U3-60, U3-72 and U3-112 have closely similar alpha subunit sequences since they differ from each other by less than 6 residues (Fig. S2). Compared to PhnI, these hybrids showed 42 or 43 amino acid replacements (Table 2), some of which are likely to perturb intramolecular interactions within alpha subunits because they are located at the interface between the catalytic and Rieske domains. These changes include Met244, which replaces a threonine in PhnI and Lys332, which replaces a serine. In addition, the sequence Val387-Gly388-Tyr389, which is located at the interface between the catalytic and Rieske domains of

adjacent alpha subunits in PhnI, is replaced by the highly charged Lys-Asp-Arg peptide in the hybrids. Such substitutions might have marked destabilizing effects on the structure of the oxygenase component. However, the U3-18 hybrid RHD displayed reasonable stability, which seems to be correlated with a unique amino acid replacement, a cysteine in position 409 (Fig S2). In the crystal structure of PhnI, a serine residue is present at this position, which is part of a α helix in the core of the catalytic domain and interacts with main chain O of Val202 and Phe242. The former residue is part of the catalytic pocket, the latter is found in a nine-stranded antiparallel β -sheet in the center of the domain. Additionally, Ser409 participates in a hydrogen bond network involving three water molecules and extending to Asn295, one of the amino acid forming the substrate-binding pocket next to the active site Fe atom (Fig. 3). This network is highly conserved across RHDs of known structure. Hence, Ser409 appears to play a critical structural role, and it is clear that a bulkier amino acid in this position would perturb interactions stabilizing the protein. In this respect, the lack of activity of the U3-60 and U3-89 hybrids might be related to the presence in position 409 of an arginine and a tryptophan, respectively. Indeed, by modeling of the serine-to-arginine substitution, we observed large perturbations at positions 203, 241 and 251 and a shift of the beta-sheet impairing interactions between the alpha and beta subunits. On the other hand, the replacement of Ser409 by a cysteine, as observed in the U3-18 and U3-116 hybrids, would not affect the inter- or intradomain interactions, inasmuch the cysteine is expected to adopt a conformation similar to the serine, and H-bonds involving the Ser O atom in PhnI would be replaced by bonds involving the Cys S atom (Fig. 3). Nevertheless, it is unclear what could explain the superior stability of the U3-18 hybrid compared to closely similar hybrids, especially those having a serine in position 409.

DISCUSSION

1 In this study, we explored the biodiversity of PAH-specific RHDs in a polluted soil by
2 combining DNA-SIP with selective PCR amplification of targeted gene sequences. Our
3 approach involved two successive screening steps, a SIP experiment allowing for the recovery
4 of genomic DNA from soil PAH degraders only, and amplification of a portion of RHD alpha
5 subunit genes encoding the catalytic domain. The set of sequences recovered in this way
6 appeared to depend on the degree of degeneracy of the primers used. With the highly
7 degenerate primer set, a significant proportion of the sequences (33%) were unrelated to
8 RHDs, but on the other hand, a greater diversity was observed among relevant sequences than
9 when a less degenerate primer set was used (Fig. 1). Most retrieved RHD sequences were
10 relatively distant from known enzymes available in the UniProt database, and were not
11 detected in previous studies targeting PAH-specific RHDs in soil (Cebron et al. 2008; Ding et
12 al. 2010) or marine environment (Lozada et al. 2008). The predominance of unknown RHD
13 sequences is consistent with the finding that the majority of bacteria identified as
14 phenanthrene degraders in the same soil sample were either from uncultured genera or
15 unknown for their ability to degrade PAHs (Martin et al. 2012).

16 We then searched for correlations between the expected bacterial hosts of RHDs found
17 in this study, and the taxa previously identified as the main phenanthrene degraders based on
18 16S rRNA sequence analysis. Among Alphaproteobacteria, Sphingomonads were the most
19 abundant degraders, consistent with the occurrence of 11 RHD sequences affiliated to this
20 taxon (cluster 1), representing 22% of the sequences in the DcatdioxA library. Cluster 2
21 sequences appeared to be best related to RHDs from Alphaproteobacteria, although the
22 similarity level was relatively low, so their affiliation to another taxonomic group such as the
23 *Cycloclasticus* genus cannot be ruled out. Cluster 3 sequences are similar to 3-component
24 salicylate hydroxylases, which play a role in the lower part of the PAH degradation pathway
25 (Jouanneau et al. 2007). Since the latter sequences were not included as targets for PCR

1 primer design, their retrieval was unexpected. Analysis of the cluster 3 sequences indicates
2 that the reverse primer does not match the 3' end. Nevertheless, the high degeneracy of the
3 primer allowed annealing within the 3' end of the gene sequences, thus generating products
4 about 100-bp shorter than sequences in other clusters. Interestingly, the genomes of
5 Sphingomonads were found to contain multiple copies of highly similar genes related to
6 cluster 3 sequences (Pinyakong et al. 2003a; Schuler et al. 2009), suggesting that they might
7 be present in higher copy numbers than other catabolic genes in metagenomic DNA from the
8 studied soil. This might account for the relatively high proportion of cluster 3 sequences
9 (30%) in the Dcatdiox A library.

10 Using a second primer pair targeting Betaproteobacteria RHDs, sequences closely
11 related to *Acidovorax* RHD were obtained, consistent with the prevalence of this genus
12 among the soil phenanthrene degraders (Martin et al. 2012; Singleton et al. 2005). However,
13 the majority of the sequences in the DcatdioxB library (87%) showed no clear affiliation.
14 Since uncultured members of the Rhodocyclaceae also appeared as prominent degraders in
15 our previous SIP experiments, we suggest that the sequences of interest (cluster 4) might be
16 specific of this bacterial taxon. Rhodocyclaceae have been detected as potent PAH degraders
17 in other SIP studies (Singleton et al. 2007; Singleton et al. 2006), but no representative strain
18 has been isolated so far. Therefore, no RHD sequence specific for this taxon is yet available.

19 In a second part of this study, we demonstrated the feasibility of constructing RHD
20 chimeras to assess the catalytic activity and substrate specificity associated to partial gene
21 sequences retrieved from soil. Three RHD hybrids were proved competent in the
22 dioxygenation of 2- and 3-ring PAHs, whereas most other hybrids appeared unstable. The
23 reason for this instability is intriguing. Four of them have very similar sequences but variable
24 stabilities. The U3-18, U3-60, U3-72 and U3-112 sequences are almost identical to the
25 catalytic domain of strain LH128 RHD (Schuler et al. 2009). Mismatches with the LH128

sequence occur at positions where amino acids were neither strictly conserved nor involved in catalytic activity, namely, Val182, Lys195, Gln273, Asn370, Ser409 (Fig. S2). Therefore, the question arises as to why RHD chimeras resulting from a sequence combination between the α subunits of two naturally occurring enzymes were so unstable. In this respect, the case of the U3-18 hybrid is interesting in that it showed reasonable stability and activity, and displayed one unique replacement compared to the other three hybrids in position 409. A cysteine is found at this position whereas an arginine occurs in U3-60 and a serine in the other hybrids as well as in most RHD sequences in the databases (Fig. S2). This substitution appeared to be the sole sequence difference between the U3-60 and U3-18 hybrids, and therefore this unique change has drastic effects on the enzyme stability. Structural considerations could predict the strong destabilizing effect of a bulky residue such as arginine in position 409 on the 3D fold of the RHD catalytic component and thus explain the instability of the U3-60 hybrid. On the other hand, the instability of other hybrids with a serine in position 409 could not be interpreted on structural grounds.

During the course of this study, Standfuß-Gabisch reported on the characterization of hybrid biphenyl dioxygenases generated through a similar strategy (Standfuss-Gabisch et al. 2012). Sequences amplified from soil DNA encoded a portion of the catalytic domain similar to that considered in the present work, but diversity was lower as most sequences fell into 2 main clusters and shared 97-100% identities within clusters and 85-88% identities between clusters. Hybrid RHDs were constructed using the well-characterized biphenyl dioxygenase from *B. xenovorans* LB400. Out of 21 hybrid RHDs tested, three were found to be unstable and three were inactive. No obvious correlation was observed between lack of activity and predicted structural changes resulting from amino acid substitutions. The high proportion of active hybrids might be explained by the fact that, in their case, a majority of the hybrids exhibited a catalytic domain closely similar (85-96% identity) to that of the LB400 enzyme.

1 In comparison, our constructions carried an inserted domain sharing less than 83% identity
2 with the corresponding region of the PhnI sequence. Hence, the stability and activity of
3 hybrid RHDs might depend on the relatedness between the sequences to be inserted and that
4 of the recipient enzyme system. Also, for the recovery of active hybrids, one should consider
5 the intrinsic stability of the enzyme system employed to generate hybrids or its propensity to
6 tolerate replacement of a large portion of its catalytic domain. In this respect, RHDs other
7 than PhnI from strain CHY-1 might prove more suitable for the construction of hybrid
8 enzymes with appropriate stability. Finding such enzymes is likely to be a prerequisite to the
9 functional exploration of RHD diversity in soil, based on the exchange of catalytic domains.

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17 sediments. *FEMS Microbiol Lett* 262:148-157

1 Tables

2 Table 1: Bacterial strains and some of the plasmids used in this study

| Strain or Plasmid | Description/ genotype | Reference or source |
|---------------------------------|--|------------------------------|
| <i>Escherichia coli</i> Strains | | |
| NEB 5- α | <i>fhuA2</i> Δ (<i>argF-lacZ</i>) <i>U169 phoA glnV44</i> Φ 80 Δ (<i>lacZ</i>)M15 <i>gyrA96 recA1 relA1 endA1 thi-1 hsdR17</i> | New England Biolabs |
| JM109 | <i>recA1 supE44 endA1 hsdR17 gyrA96 relA1 thi-1</i> Δ (<i>lac-proAB</i>) (Yanisch-Perron et al. 1985) F(<i>TraD36 proAB⁺ lacI^f lacZ</i> Δ M15) | |
| Plasmids ^a | | |
| pSD8 | pET15b carrying <i>phnA1aA2a</i> from strain CHY-1 | (Demaneche et al. 2004) |
| pIZ1036 | Km ^R , broad range expression plasmid derived from pBR1MCS-2 | (Moreno-Ruiz et al. 2003) |
| pIBA34 | pIZ1036 carrying <i>phnA3</i> and <i>phnA4</i> from strain CHY-1 | Jouanneau, unpublished |
| pUC18 | Amp ^R , cloning plasmid | (Yanisch-Perron et al. 1985) |
| pFMA1 | pUC18 carrying <i>phnA1aA2a</i> | This study |

3 ^a Other plasmids constructed in this study are listed in Table 2

Table 2 : Characteristics of chimerical RHDs and plasmids used for heterologous expression

| Sequence | Insert length (bp) | No of aa substitutions in the exchanged domain ^a | Expression plasmid | RHD activity ^b (%) |
|----------|-----------------------|---|-----------------------|----------------------------------|
| U3-116 | 695 | 39 | pFMA2 | 49 |
| U3-60 | 695 | 42 | pFMA3 | nd |
| U3-72 | 695 | 42 | pFMA4 | 0.26 |
| U3-89 | 695 | 104 | pFMA5 | nd |
| U3-112 | 695 | 43 | pFMA12 | nd |
| U3-18 | 695 | 43 | pFMA13 | 34 |
| U3-16 | 580 | 133 | pFMA6 | nd |
| U3-124 | 580 | 133 | pFMA7 | nd |

^a Numbers refer to amino acid changes compared to the corresponding region of PhnA1a

^b Activities were determined with naphthalene as substrate and calculated as percentages of the activity measured with the reference strain carrying pFMA1. nd means not detected.

Table 3 : Relative activity of hybrid dioxygenases with 2- and 3-ring PAHs

| PAH Hybrid | Naphthalene | Biphenyle | | Phenanthrene | | Anthracene | |
|---------------|--------------------|------------|----------------|--------------|----|------------|-----|
| | mU/ml ^a | mU/ml | % ^b | mU/ml | % | mU/ml | % |
| PhnI | 3020 (230) | 840 (30) | 28 | 1300 (260) | 43 | 295 (20) | 9.4 |
| U3-116 | 1480 (60) | 350 (3) | 24 | 550 (25) | 37 | 70 (1) | 4.8 |
| U3-18 | 1030 (20) | 150 (40) | 15 | 447 (30) | 43 | 21.8 (1) | 2.1 |
| U3-72 | 7.83 (1) | 3.68 (0.6) | 47 | 5.25 (0.8) | 67 | 1.73(0.16) | 22 |

^a One milliunit of activity (mU) is the amount of enzyme that produced one pmol of dihydrodiol per hour at 25°C under the defined assay conditions. Numbers in parenthesis are standard deviations.

^b Percentages were calculated relative to the activity measured with naphthalene as substrate.

Figure legends

Figure 1 : Phylogenetic tree of predicted partial sequences of RHD alpha subunits from soil subjected to DNA-SIP. Sequences were from two clone libraries obtained by PCR amplification of target RHD genes using labeled DNA extracted from soil treated with ^{13}C -phenanthrene for 5 days. Alignment was performed on a common portion of the translated sequences (corresponding to residues 98-411 of PhnA1a) encompassing the catalytic domain of RHD α -subunits. Only 47 of the 50 RHD sequences of the DcatdioxA library were considered in the alignment. Five clusters of related sequences were defined (see text). Triangles represent groups of sequences sharing >97% identity, with sequence number per group indicated on the left. Arrows designate sequences used for the construction of hybrid RHDs. Two labels present at the end of a branch (i.e. U3_72 U3_6) means that two identical sequences were found at this position. Relevant accession numbers are indicated.

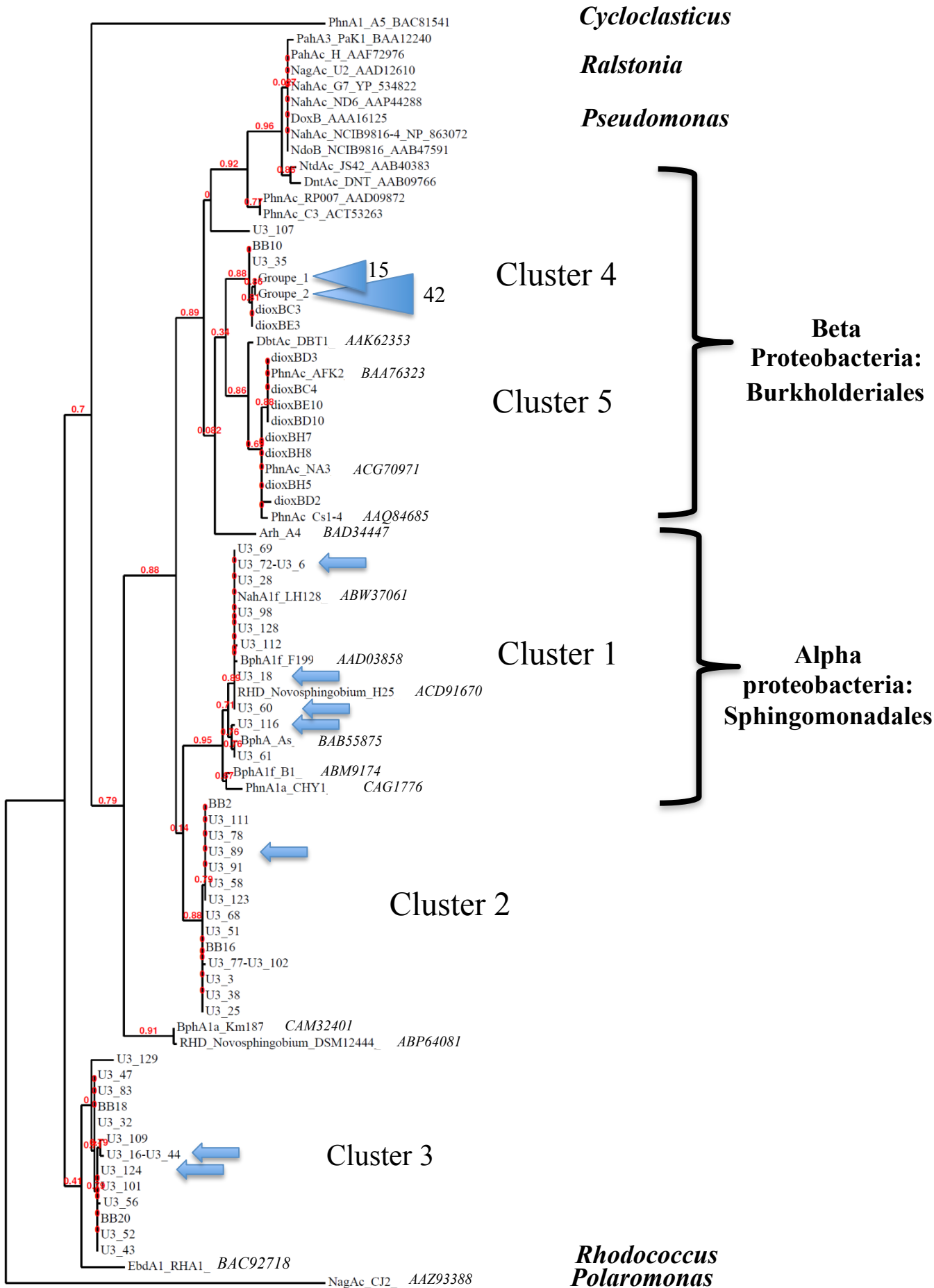
Figure 2 : Immunoblot analysis of protein cell extracts from strain JM109 expressing hybrid RHDs.

Panel A: whole cell extracts prepared by lysing cells in SDS mix 10 min at 99°C. Protein loading in each lane was adjusted based on equal bacterial density (OD_{600}) as measured prior to cell lysis. Lane 1 : non-induced cells carrying pFMA13 (U3-18); lane 8: purified ht-PhnI, 0.2 μg . Panel B: Cytosolic fractions prepared by cell ultrasonication and centrifugation. Lanes were loaded with 6 μg (lanes 2-4) or 12 μg (lanes 5-7) of proteins. Lanes 1 and 8 were loaded with 0.3 and 0.1 μg of purified ht-PhnI, respectively. Other lanes were loaded with extracts from cells expressing PhnI (lane 2) or the following hybrid RHDs; lane 3, U3-116 ; lane 4, U3-18 ; lane 5, U3-60 ; lane 6, U3-72 ; lane 7, U3-112. Western blots were revealed

1 with anti-Histag antibodies. The second band detected with $M_r \approx 45,000$ is the reductase ht-
2 PhnA4 co-expressed in JM109.

3
4 Figure 3 : Close up view of the hydrogen bond network involving Ser409 and its connections
5 with residues at the active site of Phn1.

6 Ser409 is involved in a hydrogen bond network (red dashed lines) involving two water
7 molecules, and extending to Asn295, a residue lining the catalytic pocket. It is connected
8 through its hydroxyl group to the carbonyl of Val202 and Phe242. The network also includes
9 the Gly203-Asp204 dipeptide, where Asp204 is thought to be responsible for electron transfer
10 between the Rieske center of the adjacent alpha subunit and the catalytic site. The covalent
11 bonds between active site Fe atom and its three ligands are shown as green dashed lines. An
12 indole molecule is represented inside the substrate-binding site. The figure was prepared with
13 Molscript (Kraulis 1991) and Raster3D (Merritt and Murphy 1994).



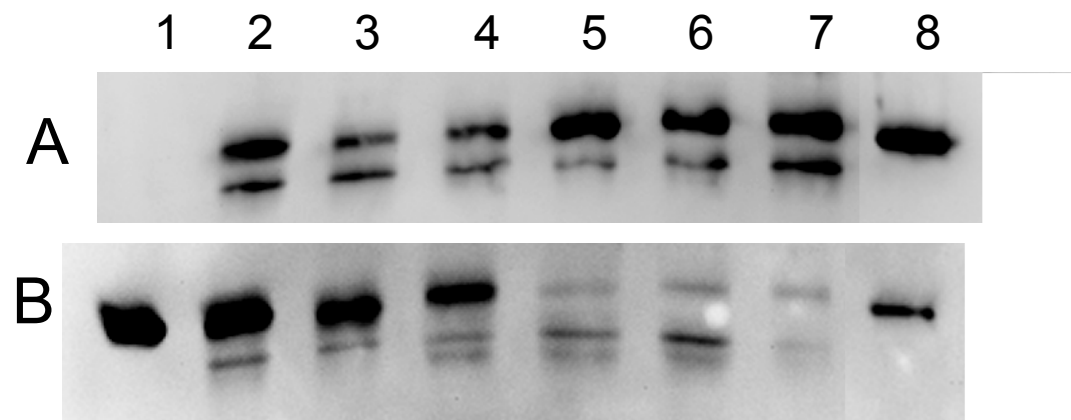


Fig. 2: Martin et al

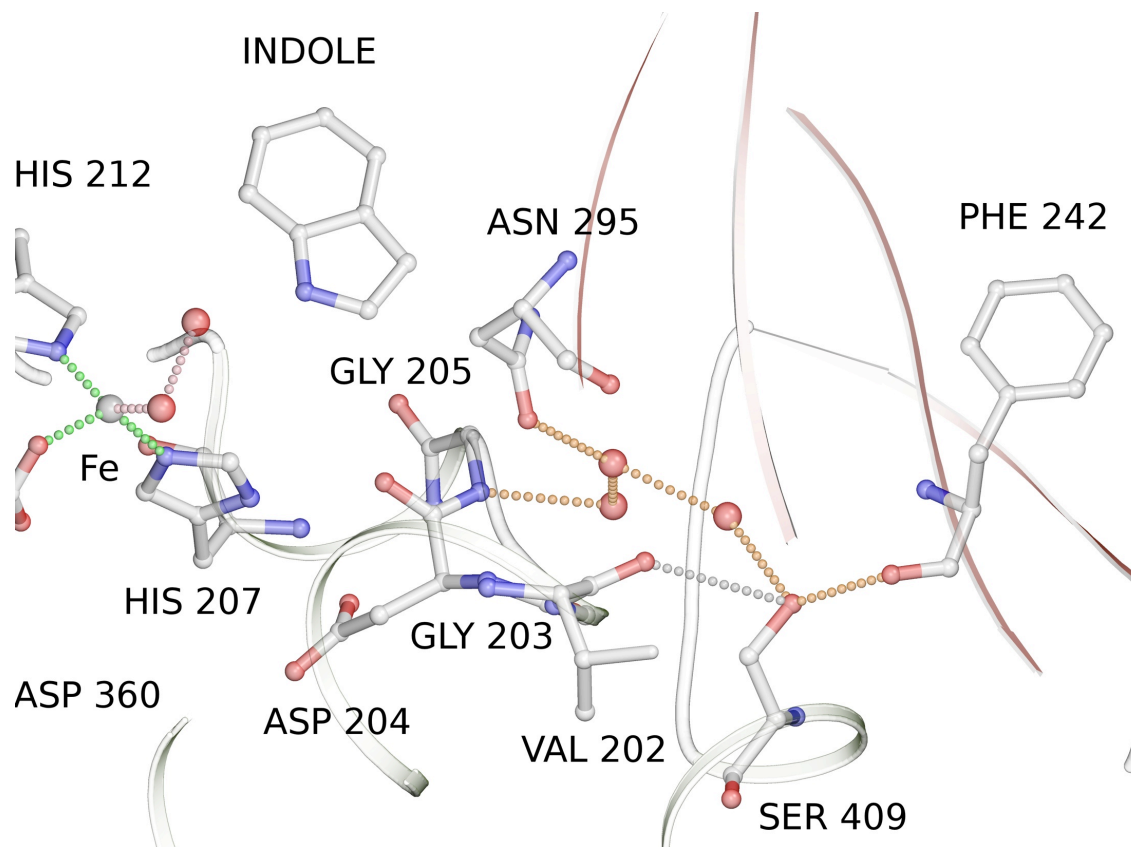


Fig. 3: Martin et al.